
(12) UK Patent Application (19) GB (11) 2 013 691 A

(21) Application No 7903836

(22) Date of filing 2 Feb 1979

(23) Claims filed 2 Feb 1979

(30) Priority data

(31) 875140

(32) 6 Feb 1978

(33) United States of America
(US)

(43) Application published
15 Aug 1979

(51) INT CL²

C07G 7/00 A61K 37/06
(A61K 39/02 39/40)

(52) Domestic classification

C3H FX

A5B 102 105 132 133

134 137 A

(56) Documents cited

GB 1211876

(58) Field of search

A5B

C3H

(71) Applicant

Stolle Research and
Development

Corporation, 6990 Cornell

Road, Cincinnati, Ohio

45242, United States of

America

(72) Inventors

Ralph J. Stolle,

Lee R. Beck

(74) Agent

Carpmaels & Ransford

(54) IgG Preparations

(57) There is disclosed an immune
gammaglobulin (IgG) preparation for
the treatment and prevention of
rheumatoid arthritis. The treatment
involves passive immunization against
a mixed spectrum of infectious
bacteria which reside in the human
gastrointestinal tract. The passive
immunization may be accomplished

by oral ingestion of IgG
immunoglobulin obtained from the
milk of cows that have been
immunized against a specific
spectrum of bacterial types. A unique
combination of bacterial species is
formulated into a vaccine which may
then be used to immunize dairy cattle.
Preferably, the IgG preparation is
obtained from the milk of the
immunized cows.

GB 2 013 691 A

2013091

1 / 3

FIG. 1

Monthly Questionnaire and Scoring Guide

Date _____

Please answer the questions by filling in the blank spaces or checking the boxes.

Name _____ Sex _____ Age _____

Race _____ Marital Status: ☐ Married ☐ Unmarried ☐ Widowed
Employment: ☐ Full-time ☐ Part-time

How long have you had arthritis? _____ years

Score 1. This morning, did your stiffness last:0 longer than 30 minutes
or
1 less than 30 minutes2. This question is about your joint pains in just this past week only:

JOINTS	NO PAIN	PAIN LASTING ONE DAY OR LESS	PAIN LASTING CONSTANTLY FOR MORE THAN ONE DAY
	Score 0	Score 1	Score 2
a. Shoulders			
b. Elbows			
c. Wrists			
d. Hands			
e. Hips			
f. Knees			
g. Ankles			
h. Feet			

3. Please tell us the drugs you took yesterday: (Pills)

Score #Pills	a. Aspirin (any form, Ecotrin, Bufferin, Anacin, etc.)	<input type="radio"/> Yes <input type="radio"/> No	How many yesterday? _____
#mgx4	b. Cortisone (any form)	<input type="radio"/> Yes <input type="radio"/> No	How many yesterday? _____
#Pills x 2.5	c. Indocin (blue & white capsules)	<input type="radio"/> Yes <input type="radio"/> No	How many yesterday? _____
#grx2	d. Pain Pills (Darvon, Codein etc.)	<input type="radio"/> Yes <input type="radio"/> No	How many yesterday? _____
#Pills x 7	e. Butazolidin	<input type="radio"/> Yes <input type="radio"/> No	How many yesterday? _____

FIG. 1A

4. In the last 3 months, have you had: (Other medication)

Score 1	Gold	<input type="radio"/> Yes
Score 2		<input type="radio"/> No
Score 1	Plaquenil	<input type="radio"/> Yes
Score 2		<input type="radio"/> No
Score 1	Cortisone Shots	<input type="radio"/> Yes
		<input type="radio"/> No

5. In the past month, are you: (ADL)

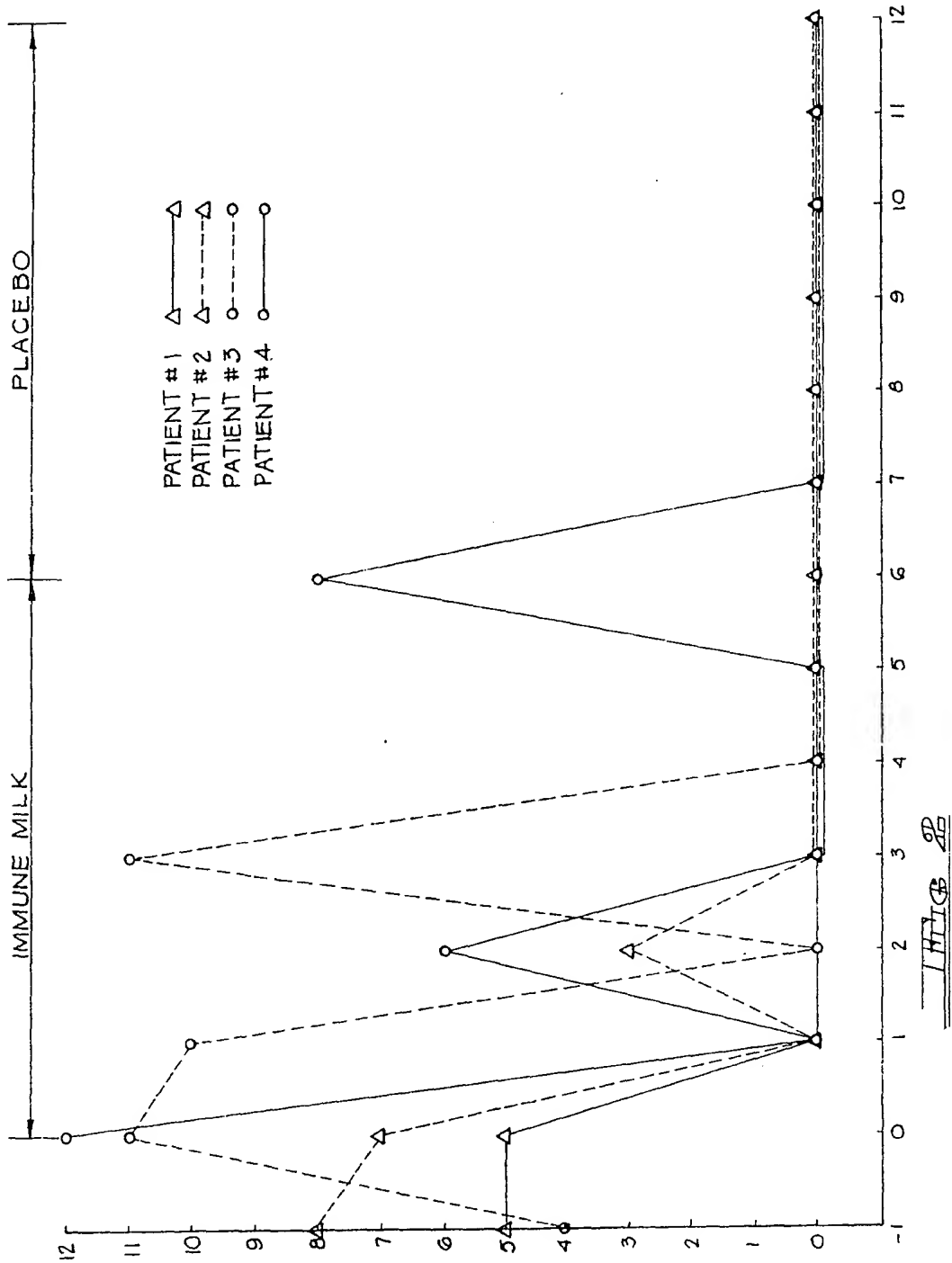
<u>Score</u>	
1	<input type="radio"/> Able to carry out all normal activities, (work, housework, shopping)
2	<input type="radio"/> Able to carry out all normal activities but with some limitations (limited housework, limited shopping, etc.)
3	<input type="radio"/> Able to carry out only some of your normal activities because of joint problem
4	<input type="radio"/> Are you able to carry out only <u>a few</u> of your normal activities
5	<input type="radio"/> Are you very dependent on others for your own care
6	<input type="radio"/> Unable to get out of chair or bed by yourself

6. Tell us how your arthritis is bothering you. (Monthly change)

<u>Score</u>	
1	a. Joint Pain: <input type="radio"/> worse than last month
2	<input type="radio"/> same as last month
3	<input type="radio"/> less than last month
1	b. Morning Stiffness: <input type="radio"/> longer than last month
2	<input type="radio"/> same as last month
3	<input type="radio"/> shorter than last month
1	c. Joint Swellings: <input type="radio"/> worse
2	<input type="radio"/> same
3	<input type="radio"/> less

2013091

3 / 3



et al, 1970), and very little IgG finds its way into the gastrointestinal fluids. Another important difference between the classes of immunoglobulin is related to their metabolic rate. The degradation of each class of immunoglobulin, regardless of its location within the body, appears to be under separate control. The functional catabolic rate varies from as low as 6.5% for IgG to as high as 90% for IgE with other classes of immunoglobulin falling in between (Waldman et al, 1970). Further, the different immunoglobulin classes also differ in their avidity with which they bind to antigens, and in their ability to combine with complement, which is one of the requisites for killing living bacterial cells (Heremans, 1960). It is important to emphasize these differences in the types of antibodies because immune effects may vary depending on the type of antibody involved.

The most commonly held theory is that the different classes of immunoglobulin have evolved to function in different environments within the body. It is known, for example, that a special and distinct immune system exists for the production of antibodies which function in the environment of the gut. Moreover, there is general agreement that the immune functions of the gut are controlled specifically by IgA antibodies and not IgG. Therefore, under natural conditions, IgA is the class of immunoglobulin which regulates immune control over bacterial infections which occur in the gastrointestinal cavity of man. Since IgG, IgM, IgD, and IgE are not normally found in the intestinal secretions, it is not logical to expect that any of these types of antibodies would be effective in treating infections in the environment of the gut.

The principal immunoglobulin in the milk of cows is IgG, not IgA (Sullivan, et al, 1969). Therefore, bovine milk is not an obvious source of antibody for treating bacterial infections of the gut in man because of its high concentrations of IgG and low concentrations of IgA.

The method of immunization is another important parameter when considering the different classes of immunoglobulin. It is well known to those skilled in the art that different methods of immunization result in the preferential production of different types of antibodies. For example, local immunization of secretory tissues achieved by exposing the tissue of antigens stimulates the preferential production and secretion of IgA type immunoglobulins. The technique of intramammary perfusion as described in the Petersen patent (U.S. Patent 3,376,198) is an example of local immunization. This method stimulates production and secretion of IgA antibodies and is not an effective method for producing IgG.

To produce the preparation of the present invention, intramuscular injection is preferably used, because IgG is the principal immunoglobulin in cow's milk, not IgA, and in the cow, systemic immunization is the preferred method for generating IgG type antibodies in milk. This distinction between the IGG and IGA type immunoglobulin is important because it teaches that systemic immunization and not local immunization is the preferred method for obtaining milk antibodies of high titer. Moreover, this distinction teaches that the immune products produced by mammary perfusion of a vaccine are distinctly different from the immune product produced by intramuscular injection of the identical vaccine. Thus, the product of this invention (IgG antibodies) is distinctly different from the product obtained by the Petersen process.

The immune product of this invention is an improvement over the product of Petersen's invention because the concentration of antibodies of the IgG type is significantly higher than the concentration of antibodies of the IgA type. There is no evidence in the literature to support the claim that IgG antibodies can be produced by intramammary perfusion of antigens. Moreover, since the levels of IgA immunoglobulins are either non-existent or extremely low in cow's milk, it is unreasonable to suggest that the teaching of Petersen's patent has any relevance to the claim of this invention. On the contrary, the teaching of the Petersen patent leads away from the discovery of this invention since it implies that IgA is a biologically active factor in cow's milk, which has potential therapeutic application.

Thus, the preparation of the present invention is preferably obtained by formulating a unique combination of bacterial species into a vaccine, which is administered to healthy dairy cows. The IgG antibodies obtained from the milk of the immunized cows constitute the preparation of the invention which may be used in the method of treatment involving the passive immunization of the patient by oral ingestion of the IgG immunoglobulin. This passively immunizes against a mixed spectrum of infectious bacteria which reside in the gastro-intestinal tract. This treatment eliminates conditions in the gastro-intestinal tract which cause rheumatoid arthritis.

The invention will now be described with reference to the accompanying drawings, in which:—

Figure 1 is a specimen of a questionnaire referred to in the specification;

Figure 1a is a continuation of the questionnaire of Figure 1, and

Figure 2 is a graph plotting results of test in-terms of RF titer against time, over a 12 month period, 6 months on immune milk and 6 months on placebo.

In a preferred embodiment, preparation of this invention is a low-fat powdered milk which contains a population of natural IgG type antibodies that react with the bacterial species listed in Table 1.

Table 1
Bacterial Antigens

	Organism	*ATCC NO.	
	<i>Staphylococcus aureus</i>	11631	
5	<i>Staphylococcus epidermidis</i>	155	5
	<i>Streptococcus pyogenes</i> , A. Type 1	8671	
	<i>Streptococcus pyogenes</i> , Type 3	10389	
	<i>Streptococcus pyogenes</i> , Type 5	12347	
	<i>Streptococcus pyogenes</i> , Type 8	12349	
10	<i>Streptococcus pyogenes</i> , Type 12	11434	10
	<i>Streptococcus pyogenes</i> , Type 14	12972	
	<i>Streptococcus pyogenes</i> , Type 18	12357	
	<i>Streptococcus pyogenes</i> , Type 22	10403	
	<i>Aerobacter aerogenes</i>	884	
15	<i>Escherichia coli</i>	26	15
	<i>Salmonella enteritidis</i>	13076	
	<i>Pseudomonas aeruginosa</i>	7700	
	<i>Klebsiella pneumoniae</i>	9590	
	<i>Salmonella typhimurium</i>	13311	
20	<i>Haemophilus influenzae</i>	9333	20
	<i>Streptococcus viridans</i>	6249	
	<i>Proteus Vulgaris</i>	13315	
	<i>Shigella dysenteriae</i>	11835	
25	<i>Streptococcus</i> , Group B		25
	<i>Diplococcus pneumoniae</i>		
	<i>Streptococcus mutans</i>		

Corynebacterium. Acne, Types 1 & 2

American Type Culture Collection, 12301 Parklawn Dr., Rockville, Md. 20852

- 30 The antibacterial milk contains all of the substances normally found in low-fat powdered milk.
The principal constituents comprising antibacterial milk are shown in Table 2. 30

Table 2
Quantitative and Qualitative Analysis of Antibacterial Milk

	Proteins	35.6%	
	Fat	1.0%	
35	Carbohydrates	52%	35
	Minerals	7.8%	
	Moisture	3.5%	

Each reliquified quart of 3—4 ounces of non-fat dry milk contains approximately:

40	1200 mg calcium	157%	40
	935 mg phosphorous	125%	
	0.3 mg thiamine	32%	
	1.78 mg riboflavin	140%	
	1.04 mg niacin	10%	
	324 Calories		

- 45 Antibacterial milk and normal cow's milk contain the same approximate percent by weight concentration of ingredients. Moreover, the concentration of type IgG immunoglobulin in antibacterial milk and normal milk is identical. Therefore, it is only the specificity of antibodies comprising the antibacterial milk which distinguishes it from normal milk. By specificity of the immunoglobulin is meant the spectrum of bacterial species that the antibodies react with. 45

- 50 Antibacterial milk contains no drug additives or any other components which are not natural food products of the cow. 50

The immune milk, which comprises the preferred embodiment of the present invention, is also useful in the control of auto-immune diseases, e.g. lupus erythematosus, which are caused or aggravated by bacterial inventions in the gastrointestinal tract.

- 55 The polyvalent antigen used for the induction of the antibacterial milk may be prepared as follows:— 55

Preparation of the Vaccine

The bacterial strains listed in Table 1 were obtained from the American Type Culture Collection,

which ensures authenticity of bacterial strains and the highest standard of purity that is available. Upon receipt, each individual bacterial strain was grown on a blood agar plate to test the viability of the culture and to determine if growth pattern is typical or atypical of the bacteria in question. A single colony from each of the test cultures was taken for histological examination to further ensure authenticity and plurality of the culture. A single colony of each culture was used to inoculate 500 ml of standard culture broth. The standard broths recommended by the American Type Culture Collection were used to grow each of the specific bacteria listed in Table 1.

All organisms were incubated as static cultures with the exception of 12, 13, 14 and 60, which were incubated in the shaker to provide agitation. Identification of bacterial strains and the American Type Culture Collection catalog numbers are shown in Table 1. Each culture was cultivated for 48 hours at 37°C. Following incubation, the cultures were killed by heating at 60°C for two hours. Samples of the killed bacteria were used to inoculate fresh broth which was then incubated for 24 hours at 37°C to determine if the killing process was complete. Only cultures proven sterile by this procedure were used for further processing. Sterile cultures were then washed five times in distilled water and the cells were recovered by centrifugation. The bacterial cells were frozen by immersion in liquid nitrogen and freeze-dried by the process of lyophilization. The lyophilized cells were stored in sterile vials until used for production of the polyvalent vaccine. The polyvalent vaccine was prepared by weighing out one gram quantities of each of the bacterial strains. The dry cells were mixed together and this mixture was suspended in sterile physiological saline (20 grams of bacteria per 500 ml saline).

A sample of the concentrated solution was diluted in serial fashion with saline to determine dilution which gives a concentration of 4×10^8 ml per cc. The stock concentrated polyvalent vaccine was dispersed into multiple containers and stored frozen. A sufficient amount of concentrated antigen was included in each individual container to immunize 50 cows. The final dilution of concentrate was made just prior to immunization. The preferred procedure is to remove a sufficient number of vials to immunize the number of cows to be treated. For example, the vials are removed 24 hours prior to the planned time of immunization; a sample of the concentrate is then diluted in a sterile container to a final concentration of 4×10^8 cells per ml. The maximum response in cows is obtained by injecting 20×10^8 bacterial cells or 5 cc of the sterile preparation which is 4×10^8 cells per ml according to the method of immunization described below.

30 Preferred Process for Immunization of Cows

The antibody product of the invention is produced by immunizing cows with the polyvalent antigen prepared as described above. The cows are injected with 5 cc of polyvalent antigen containing 20×10^8 bacterial cells. The injection is made intramuscularly in the gluteus maximus muscle of the hind leg. This procedure is repeated at one week intervals for four consecutive weeks beginning 2—3 weeks prior to the predicted day of parturition. Following the primary immunization, booster injections using the same concentration of the antigen, are given every 14 days. This method of immunization gives the maximum antibody titer.

Preferred Collection, Handling and Processing of Milk

The milk is collected from immunized cows in a modern dairy parlor. A fully automated milking system collects and stores the milk under complete sanitary conditions. The milking system consists of automated machines connected directly to refrigerated storage tanks by a closed system of pipes. The complete system is cleaned and sterilized following each milking to ensure maximum sanitary conditions. It is important to take careful steps to prevent the growth of bacteria to immune milk during processing, since such bacteria can lower the titer of antibodies in the milk.

Milk is transported daily from the refrigerated holding tanks to a dairy processing plant by milk transport trucks. At the dairy plant a high temperature short-time system is used to pasteurize the antibacterial milk. Specialized dairy machinery provides the flash heating of a continuous flow of milk to 155°F for a period of not more than 15 seconds. Temperature and time is critical since antibody is susceptible to degradation by heat. Milk antibody is destroyed at temperatures above 165°F, if held for periods longer than one minute. Following pasteurization, the whole milk is immediately cooled and the fat is removed by centrifugation, and the skimmed whole antibacterial milk is powdered by a spray process. The spray process consists of a large drying chamber into which hot air (350°F) is blown at high velocity. The skimmed milk is atomized into the chamber where the finely divided milk particles are instantly dried as they fall to the bottom of the tank. The dried milk is removed automatically by means of mechanical devices and the milk powder is packaged under sanitary conditions. Prior to atomizing, the skimmed milk is condensed by boiling in a chamber under vacuum (100—110°F). At each step it is critical to keep the bacteria from contaminating the milk since this reduces the titer of the antibody.

Preferred Testing Procedures

Immune milk was prepared in inbred Holstein cows. The cows were immunized by the intramuscular injection of a mixture of bacterial antigens identified in Table 1. The vaccine was prepared by the process described above. The immunologic response of the cows was boosted by bi-

weekly injections of the vaccine. The milk from these cows was pooled, the fat removed, and the non-fat milk was pasteurized by exposure to 160°F for 16 seconds followed by a spray-drying process in which the temperature of the milk did not exceed 85°F. The milk was packaged in one quart polyethylene containers. Control milk (placebo) was non-fat powdered milk purchased from a local producer.

Erythrocyte sedimentation rates were determined on freshly collected blood by the method of Westergren and corrected for hematocrit according to the method of Wintrobe & Langsberg (1935). Rheumatoid factor titers were determined by the Singer-Plotz (1966) macroscopic tube test.

Patients were accepted for the study on the basis of an elevated erythrocyte sedimentation rate and a positive rheumatoid factor titer. Nine patients were studied for 12 months and 11 patients were studied 18 months. The patient group was composed of thirteen caucasian females ranging in age from 32 to 69 years with an average of 50.4 years, and seven caucasian males ranging in age from 43 to 70 years with an average age of 58.1 years. The mean duration of arthritis was 10.8 years for the females and 11.0 for the males. Patients were randomly placed either on immune milk or on non-immune milk (a commercial product purchased in the Dayton area that served as a placebo). Both milk products were packaged in identical containers and were identified as being immune milk or placebo, respectively, by a blue or red pressure-sensitive label that was attached to each container at the time it was filled. The labels were removed just prior to dispensing the milk to the patients. Thus, at no time did the patients know whether they were receiving immune milk or placebo. Patients were randomly (as determined by the flip of a coin) selected to receive either immune milk or the placebo during the first six-month period. At the end of this time, those that were receiving immune milk were placed on the placebo and those that were receiving placebo were placed on immune milk for the second six-month period.

At the end of the second six-month period, 11 patients volunteered to remain on the study for an additional six months. The type of milk (immune or placebo) was again changed at this time and observations were continued. Thus, the study was comprised of three six-month periods, 11 of the patients participating for three periods and nine participating for two periods.

Patients were seen at monthly intervals at which time a one month supply of milk was dispensed, an evaluation questionnaire was filled out and a blood sample was collected for rheumatoid factor titer, erythrocyte sedimentation rate and hematocrit determination.

Patients were instructed to take a quantity of non-fat milk solids equivalent to one quart of milk post prandially two times daily. The milk solids were freshly dissolved in one pint of cool tap water immediately before ingestion shortly after awakening in the morning and again just prior to retiring at night. They were told to see their physician as usual and to follow the treatment regimens prescribed by him. Medication was to be taken ad libitum or as prescribed by their regular doctor. We requested only that they report the quantity of medicines taken.

A questionnaire was completed by each patient at monthly intervals. It was divided into six sections that deal with:

- 1) duration of morning stiffness,
 - 2) severity of pain experienced in each of eight joints,
 - 3) type and quantity of drugs with short-term actions that were taken,
 - 4) type and quantity of drugs with long-lasting actions that were taken,
 - 5) ability of patient to conduct his normal activities,
- and
- 6) severity of symptoms of rheumatoid arthritis.

The numbers shown in the spaces next to each answer indicate the score assigned to that answer in the course of evaluating the questionnaires. In scoring the sections dealing with medications, an effort was made to reflect the relative anti-inflammatory and analgesic activities of the various drugs used. A five-grain aspirin tablet was assigned a value of one. All other drugs (with the exception of gold, plaquenil and cortisone shots which were considered separately) were arbitrarily assigned values relative to aspirin. Thus, all salicylate preparations, Tylenol, Darvon, Motrin, etc. were considered equivalent to a five-grain aspirin tablet and were also assigned a value of one. The number mg of Prednisone was multiplied times four, the number of Indocin capsules taken was multiplied times 2.5. The number of grains of codeine was multiplied times two, and the number of Butazoladin tablets taken was multiplied times seven.

The mean scores in each category were calculated for each six-month period. The differences of the means were then calculated by subtracting the mean values scored during administration of immune milk from those scored during administration of placebo. When the results were calculated in this manner, improvement in the patient's condition during the period he received immune milk was indicated by negative values for questions one and six, and by positive values for all other questions. Mean corrected erythrocyte sedimentation rates (ESR) and rheumatoid factor titers (RF) were respectively shown in a similar manner. These were calculated in such a way that positive values reflect a lower erythrocyte sedimentation rate or rheumatoid factor titer during administration of immune milk. The data were statistically evaluated using the Statistical Analysis System of Goodnight

et al. (SAS Institute, Raleigh, N. C.). Calculations were performed with the aid of an IBM model 370/155 computer.

Results

The immune milk was well tolerated by all patients with the exception of one who had pernicious anemia. This patient complained of diarrhea and was terminated from the study. Some patients reported a weight gain during the course of the study. This may have been due to the increased caloric intake from the milk or possibly reflects a generalized improvement in their physical condition.

Table 3

	Periods of Observation		Treatment Placebo		Regimen Immune		Mean Difference	P
	Control	Immune	Mean	C.V.*	Mean	C.V.*		
1. A. M. Stiffness	27	24	0.332	95.7	0.679	35.6	-0.347	0.0001
2. Joint Pain								
a. Shoulder	27	24	0.954	67.1	0.716	60.2	+0.238	0.0420
b. Elbow	27	24	0.752	83.7	0.613	65.9	+0.139	0.0511
c. Wrist	27	24	0.824	73.6	0.539	74.8	+0.285	0.0010
d. Hand	27	24	1.073	54.7	0.828	56.5	+0.245	0.0011
e. Hip	27	24	0.533	90.3	0.227	135.0	+0.306	0.0005
f. Knee	27	24	0.904	74.1	0.683	59.0	+0.221	0.0015
g. Ankle	26	22	0.7811	66.9	0.659	65.7	+0.1221	0.0127
h. Feet	26	22	0.948	63.7	0.729	50.9	+0.219	0.0010
3. Pills	27	24	20.663	104.1	16.515	101.3	+4.148	0.0405
4. Other Medication	27	24	0.325	140.7	0.244	175.6	+0.081	0.0276
5. ADL	27	24	2.224	36.1	1.874	29.1	+0.350	0.0023
6. Monthly Change								
a. Pain	27	24	1.903	21.8	2.247	14.1	-0.344	0.0042
b. Stiffness	27	24	1.985	18.8	2.254	12.4	-0.269	0.0024
c. Swelling	27	24	1.924	17.9	2.117	13.3	-0.193	0.00153
7. ESR	25	23	36.293	29.7	35.922	38.2	+0.371	0.7376
8. RF	27	24	6.698	45.5	6.834	41.7	-0.136	0.9635

*Coefficient of variation.

As shown in Table 3, patients were observed during a total of 27 control periods (six-month periods during which they received placebo) and 24 test periods (six-month periods during which they received immune milk). One patient had sustained a physical injury to one of his ankles and feet. The pain in these joints was not evaluated, which accounts for there being a smaller number of periods of evaluation for these joints. The erythrocyte sedimentation rates for one patient were so extremely abnormal (more than two standard deviations removed from the mean of the values for the other patients) that they were not included. This omission accounts for the smaller number of observations reported for that variable.

The mean values and coefficients of variation (C. V.) are listed in the table to reach variable. Differences between the means were calculated by subtracting the mean value obtained during the periods the patients received immune milk from that obtained during the periods they received the placebo. A favorable response to immune milk is indicated by negative values for AM stiffness (question 1) and Monthly change (questions 6a, b, and c) and by positive values for all other variables. An effective response to immune milk was obtained for all data obtained from the questionnaires. Probabilities (P) indicate a high degree of statistical significance in every instance. The small mean differences obtained for erythrocyte sedimentation rate and rheumatoid factor titer were not significant. When erythrocyte sedimentation rates were considered on an individual basis, however, four of the twenty patients studied had statistically significant decreases while receiving immune milk.

Although immune milk had no significant effect on the mean values for rheumatoid factor titer, examination of individual patients revealed some interesting responses. Seven of the twenty patients studied had negative rheumatoid factor titers on at least one occasion during the period they were receiving immune milk. Four of them became negative during the period that they received immune milk and their titers failed to become positive during the following six-month period when they received the control (placebo) milk as shown in Figure 2. Continuation of the study past this reporting period reveals that 13 of 25 patients lost the rheumatoid factor from their blood.

Discussion

The scientist in charge of this study personally interviewed each patient at monthly intervals, and

recorded their answers to the questions. Every effort was made not to influence the patient's answers. The patients were initially informed and were frequently reminded that, during certain periods of the study, they would receive a placebo. It was anticipated that this knowledge would serve as an inducement for the patients to answer the questions objectively and without bias. At no time were the patients informed whether they were receiving immune milk or the placebo.

The question regarding medication taken "yesterday" (question #3) and the question regarding gold shots, Plaquenil and cortisone, shots (question #4) are objective and are of primary importance in considering answers given to the other questions. These questions are important for two reasons:

1) if the immune milk is effective in relieving symptoms of the disease, the patient would be expected to take fewer medicines that were allowed *ad libitum*. On an average, patients reported that they took four less aspirins or their equivalent per day during the periods that they received immune milk. They also reported that they received fewer gold shots, Plaquenil and cortisone shots during these periods;

and

2) if patients took smaller quantities of analgesics and other medicines useful in the treatment of rheumatoid arthritis, one would expect them to report increased discomfort unless the immune milk was influencing the disease favourably.

As noted in Table 2, significantly less joint involvement was reported during periods that the patients received immune milk even though they were taking less medicines for their arthritis.

Patients started on the study at monthly intervals over a one-year period, and the type of milk product (immune milk or placebo) that they initially received was randomized. The observation that positive responses or improvement were obtained for all parameters of the questionnaire, and that these mean responses were statistically significant strongly indicate that immune milk had a beneficial effect on the patients. This conclusion is reinforced by the observation that 20% of the patients experiences a statistically significant ($p < 0.05$) decrease in erythrocyte sedimentation rate while receiving immune milk.

Results of the rheumatoid factor titers are difficult to evaluate. This is due to at least in part to the fact that the origin and role of rheumatoid factors in the etiology and prognosis of rheumatoid arthritis is not understood. Rose et al (1948) showed that sheep red blood cells that were sensitized with rabbit antibody underwent agglutination in the presence of blood serum from patients with rheumatoid arthritis. The test depends on the specific reaction between normal immunoglobulin (either rabbit or human IgG) with rheumatoid factors. The specificities exhibited by rheumatoid factor are like those that would be expected of antibody against IgG (Epstein et al, 1956). The presence of rheumatoid factors has been correlated with disease severity in rheumatoid arthritis and can be identified in proteins precipitated in the tissues of patients with rheumatoid arthritis. Although a small percentage of patients with rheumatoid arthritis do not have positive rheumatoid factor titers, it is generally agreed by most rheumatologists that positive agglutination reactions do not revert to negative even when the disease is in remission. De Forest et al (1958), however, described a small number of patients who had positive rheumatoid factor titers that reverted to negative following a remission. When recrudescence of the disease occurred, the test again became positive. Aho, et al (1959) noted, however, that most patients whose disease had become inactive remained serologically positive. The fact that negative titers were observed in 60% of our patients and that in half of these, the titers remained negative for six months, proves that immune milk is affecting a primary etiologic factor responsible for rheumatoid arthritis.

The effect of immune milk in alleviating the symptoms of rheumatoid arthritis is particularly relevant when considered on the basis of the recently described relationship between the histocompatibility antigens (HL—A) and the susceptibility to rheumatic disease (Brewerton, 1976). Histocompatibility antigens are genetically determined antigens that are found on all human cells. The genes controlling their inheritance are called histocompatibility genes. There are now known to be over 40 of these genetically determined antigens. They are responsible for rejection of tissue grafts made between individuals other than identical twins. Superficially the HL-A antigens resemble ABO blood groups in that they are inherited for a lifetime. Their functions is not yet known, except in the highly artificial situation produced by transplantation. It is known, however, that the histocompatibility genes are closely linked with the immune response genes on the sixth chromosome. In this relationship, they may determine the immune response of the individual to a foreign invader, such as a bacteria.

Persons with HLA—B27 appear to be particularly susceptible to a variety of rheumatic diseases. It is postulated that this histocompatibility antigen dictates a type of immune response which in the presence of other predisposing factors leads to rheumatoid arthritis. After an intestinal infection with *yersinia enterocolitica*, some patients develop an acute peripheral arthritis (Ahvonen, et al, 1969). Similarly, after *salmonella* infection, about 2% of patients develop acute peripheral arthritis (Warren, 1970). HLA—B27 was found in 43 of 49 patients with yersinia arthritis and in 15 of 16 with salmonella arthritis (Aho, 1974). It is an attractive possibility that infective agents may thrive in the intestinal tract without giving rise to local symptoms. In patients with HLA—B27, a host response is established that results in arthritis. Thus, it is not necessary for the infective agent to gain entry into the

joints. Immune milk is beneficial to patients with rheumatoid arthritis because it contains antibodies that effectively inactivate or neutralize offending bacteria and/or their metabolic products.

Claims

1. An immune gammaglobulin (1gG) preparation for the treatment of rheumatoid arthritis, said preparation being used to control mixed bacterial infections of the gastrointestinal tract. 5
2. A preparation according to claim 1, wherein the mixed bacterial infection includes two or more of the following microorganisms from American Type Culture Collection bacterial antigens:

<i>Staphylococcus aureus</i>	11631	
<i>Staphylococcus epidermidis</i>	155	
10 <i>Streptococcus pyogenes</i> , A. Type 1	8671	10
<i>Streptococcus pyogenes</i> , Type 3	10389	
<i>Streptococcus pyogenes</i> , Type 5	12347	
<i>Streptococcus pyogenes</i> , Type 8	12349	
<i>Streptococcus pyogenes</i> , Type 12	11434	
15 <i>Streptococcus pyogenes</i> , Type 14	12972	15
<i>Streptococcus pyogenes</i> , Type 18	12357	
<i>Streptococcus pyogenes</i> , Type 22	10403	
<i>Aerobacter aerogenes</i>	884	
<i>Escherichia coli</i>	26	
20 <i>Salmonella enteritidis</i>	13076	20
<i>Pseudomonas aeruginosa</i>	7700	
<i>Klebsiella pneumoniae</i>	9590	
<i>Salmonella typhimurium</i>	13311	
<i>Haemophilus influenzae</i>	9333	
25 <i>Streptococcus viridans</i>	6249	25
<i>Proteus vulgaris</i>	13315	
<i>Shigella dysenteriae</i>	11835	
<i>Streptococcus</i> , Group B		
<i>Diplococcus pneumoniae</i>		
30 <i>Streptococcus mutans</i>		30
<i>Corynebacterium</i> , <i>Acne</i> , Types 1 & 2.		
3. A preparation according to either of claims 1 or 2, said preparation being in the form of milk for oral administration.
4. A preparation according to any one of claims 1 to 3, said preparation being for oral administration and being in a vehicle that is not harmful to the gammaglobulin, said vehicle helping to prevent destruction of the gammaglobulin in the gastrointestinal tract due to the action of proteolytic enzyme and changes in pH.
5. An immune gammaglobulin preparation according to any one of claims 1 to 4, said gammaglobulin having been produced by first preparing a vaccine from killed bacteria from two or more of the American Type Culture Collection bacterial antigens including: 40

<i>Staphylococcus aureus</i>	11631	
<i>Staphylococcus epidermidis</i>	155	
<i>Streptococcus pyogenes</i> , A. Type 1	8671	
<i>Streptococcus pyogenes</i> , A. Type 3	10389	
45 <i>Streptococcus pyogenes</i> , Type 5	12347	45
<i>Streptococcus pyogenes</i> , Type 8	12349	
<i>Streptococcus pyogenes</i> , Type 12	11434	
<i>Streptococcus pyogenes</i> , Type 14	12972	
<i>Streptococcus pyogenes</i> Type 18	12357	
50 <i>Streptococcus pyogenes</i> , Type 22	10403	50
<i>Aerobacter aerogenes</i>	884	
<i>Escherichia coli</i>	26	
<i>Salmonella enteritidis</i>	13076	
<i>pseudomonas aeruginosa</i>	7700	
55 <i>Klebsiella pneumoniae</i>	9590	55
<i>Salmonella typhimurium</i>	13311	
<i>Haemophilus influenzae</i>	9333	
<i>Streptococcus viridans</i>	6249	
<i>Proteus vulgaris</i>	13315	
60 <i>Shigella dysenteriae</i>	11835	60
<i>Streptococcus</i> , Group B		
<i>Diplococcus pneumoniae</i>		
<i>Streptococcus mutans</i>		
<i>Corynebacterium</i> , <i>Acne</i> , Types 1 & 2		

- injecting said vaccine intramuscularly in healthy cows once weekly for four consecutive weeks, and twice monthly thereafter, each injection involving 20×10^8 bacterial cells; collecting the milk from the immunized cows beginning the fourth week; and testing for titer to ensure that the minimum titer against each of the bacteria is 1—500, as determined by the tube agglutination method for testing antibody titer. 5
6. A preparation according to claim 5, wherein said vaccine has been prepared by a process which includes the steps of preparing cultures of bacterial strains in appropriate buffers, heat killing the bacteria, harvesting the killed bacteria by centrifugation, washing the bacterial strains, lyophilizing said washed strains, mixing the individual bacterial types on an equal weight basis, and suspending the 10 mixed bacterial strains in a suitable vehicle for injection into cows to produce, in the milk system of the cows, an immune gammaglobulin (1gG). 10
7. A preparation according to claim 1 and substantially as hereinbefore described.

Printed for Her Majesty's Stationery Office by the Courier Press, Leamington Spa, 1979. Published by the Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from which copies may be obtained.

IgG Preparations

Patent Number: ☐ [GB2013691](#)

Publication date: 1979-08-15

Inventor(s):

Applicant(s): STOLLE RES & DEV

Requested Patent: ☐ [JP54113425](#)

Application Number: GB19790003836 19790202

Priority Number (s): US19780875140 19780206

IPC Classification: C07G7/00; A61K37/06; A61K39/02; A61K39/40

EC Classification: [A61K39/116](#), [C07K16/12](#), [A61K31/43](#), [A61K31/65](#)

Equivalents: ☐ [CH651210](#), ☐ [DE2904044](#), ☐ [DK158138B](#), [DK158138C](#), [DK49479](#),
☐ [FR2416015](#), [HK30283](#), ☐ [IT1116524](#), [JP1060455B](#), [JP1578209C](#), ☐ [NL7900766](#),
☐ [SE448344](#), [SE7900798](#)

Abstract

There is disclosed an immune gammaglobulin (IgG) preparation for the treatment and prevention of rheumatoid arthritis. The treatment involves passive immunization against a mixed spectrum of infectious bacteria which reside in the human gastrointestinal tract. The passive immunization may be accomplished by oral ingestion of IgG immunoglobulin obtained from the milk of cows that have been immunized against a specific spectrum of bacterial types. A unique combination of bacterial species is formulated into a vaccine which may then be used to immunize dairy cattle. Preferably, the IgG preparation is obtained from the milk of the immunized cows.

Data supplied from the esp@cenet database - I2